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(54) Adsorbent for separating gases

(57) An adorbert for separating passe, which comprehese a binder and cyretatine low-falls full plast layer private a binder and a cyretatine low-falls full plast layer passed with a BC/J/L/C/D, moler ratio of from 1.9 to 2.1, wherein the exercise port dismater of the memoprose is causal to or larger than the mean free path of an adsorbuide component when describing the adorbuide component from the adorbuide, and at least 70% of the total volume of the macroprose is coughed by macroprose having a dismater equal to or larger than the mean free path of the adorbuide component.

# Description

- (000). The present investion makes to an advolvent for requesting gases to separate a gas indiance having a component which is all self sections (contrable component) and a component within its all self sections (contrable component), and a process for producing in. Particularly, the advolvent for separating passes of the created invention inclusion to an advolvent for separating passes to respect as gas mixture by a reviews every advolvent invention inclusion to an advolvent for separating passes to respective to generating an extraction of the component of the component of the component of the present invention may be executed and recovered by PRA method by using the disclosed for separating gases of the present invention may be executed and recovered by PRA method by using the disclosed for separating gases of the present invention may be executed and recovered by PRA method by using the disclosed for separating gases of the present invention may be executed and recovered by the present invention may be executed and recovered by the present invention of the present invention may be executed and recovered by the present invention of the present invention may be executed and the present invention may be executed and recovered by the present invention of the present invention may be executed and the present invention of the present invention may be executed and the present invention of the present
- bon dicelets, hydrogen or celebron microardes. [5002] Among these, acygen gas is one of the particularly important inclustrial gases, and widely used for steal making, blasching of pulp or the list. Particularly, in recent years, in order to reduce generation of NCv which is linevisible by combustion in all, acygen-enrighted combustion is used predicably in the field of inhuse burning, glass meiting or the like, and copyring as is becoming forcerated in leview of predicable professionable professionable and control professionable professio
- [0003] As an includable method for producing oxyging qus. PSA method, a cryopanic delitation processing or a manybrane separation method has been incream. Among these, PSA method within its advantageous in view of the purity or oxygen gas and cost, is increasingly used. The method for producing oxygen gas by PSA is to addoct hitting on a size on an addoctor, and debat in the remaining concentrated dioxygen as a product, set the discoherent to be used, an
- civygen gas and cost, is increasingly used. The method for producing oxygen gas by PSA is to addoct infrogen gas in air own au stability and cost in the method group gas as a product, but the discribent to be used, an air own au stability and are supported by the product of t
- introgen in the Art is selectively action for one or systalline actiotie to carry out separation of the Air Genetive actions for introgen on the or systalline action (and use the strong interested between qualityel moment of introgen and electrostate brice or attraction of cestors in the action. Therefore, for PSA matted, or systalline action is serve wherein also crossate for word of attraction or cestors in the action. Therefore, for PSA matted, or systalline action is exceed wherein also crossate for word of attraction or cestors in the action is the action of the ac
- used. Particularly, filtium-exchanged crystaline zoolte X which is for-exchanged with fifthirm calons, is excellent in selective adoption of integra, and used us a crystaline zeatile to obtain concentrated copyre by PSA ineffect. [9003] For example, US-A3, 140,939 proposes ellifitum-exchanged crystaline zeatile X which is excellent in the expanding manner of difference restartional proposes and proposed control of control and control of contr
- but an example of integrated and the second of the second
- see day in the position both and reduce the load to the viscours jumps or the bower constituting the PSR appearate, the crystalline according to formed into beach or position by very lay before, a get in trappertice from a the size of or dwarrier, so cold, no register according to the purpose, in the appointments, a network of managones in so cold in a register according to the purpose, in the appointments, a network of managones in the organization according to the reduce of the appointments or compared to advance on the advances of the crystalline accold cold length or the cold or the accordinate component address in the managones and relatable to the advances that are the center of the appointments. And the advanced component defends from the and relatable to the advances that are the center of the appointments. And the advanced component defends from the contraction of the advances of the advances of the advances of the appointments of the advances of the advances
- ackerplion site diffuses in the macropores and is evacuated to the science of the agglements. In order to obtain his expected perhamance of an admixted successful size of the proposed wherein even the adequation rate addiling at the center of the agglementate can be effectively used, so that the efficiency for adecaption of the adsorbable component can be model high.
- [DOTT] For example, a south augmanests for requesting passes wherein hypos acides and separate with addition acides in a least of 23 mill (JA-A-26 19258), or a called againment wherein the series is used, and the miscropers volume as to a few and the series of the ser
- case of separating at it, additioned on the crystalline zerolle, and for reduce the time of contracting to page minture with the decident care for consideration group in the presentance when the decident decident decident confidence component in described curried minduced pressure. As althoroption and description are repeated in PSA method, to order to detail the performance of the additional, it is necessary to improve all only the definition and the additional component when additional but also the distriction relief or the distriction relief or the distriction and the contraction and additional component when decided Accordingly, rener of the convenience all contractions are not to representations and an indistriction reconstructing passes, of which the sufferior has the best multiplication plants.
- ce (0009) The shape of the adoctions for operating gases is usually cylindrical type cellet or in the form of beads. The cylindrical type pollet and the beads or proprietd usually by estimation and untraining prantitions in discontinuous prantitions of the cylindrical system of the cylindrical system

clay. Such an adsorbent has a higher resistance against gas diffusion in the inside of the applomanate, and it is not opportunity.

- sible to effectively use the center portion of the adsorbent. [0010] Adsorption of the adsorbable component on the crystalline zeolite is an exothermic process. However, decorp-
- tion of the adsorbable component from the crystalline zeolite is an endothermic process, and a higher energy is required for description of the adsorbable component than adsorption of the adsorbable component. Accordingly to obtain the desired gas separation performance, it is required to quickly evacuate the adsorbable component described from the crystalline zealite to the exterior of the advanture. Particularly in the case where the amount of the arts chable component adsorbed on the crystaline zeolite is large, in order to evacuate a larger quantity of the adsorbable compo-
- nent to the exterior of the adsorbent during description, a higher description rate is required. A high adsorption rate when so adsorbing the edsorbable component is also an important factor. Further, although the adsorbent for separating cases is used in a state where weter is removed (activated state), the crystalline zeelite has a strong affinity with water, and there is a fear that water in the etmosphere may be re-adsorbed. If water remains in the adsorbent due to water adsorbent tion, the adsorption site for gas is occupied with water, thereby ass separation deteriorates, and the desired perform-
- ance is less likely to be obtained. 15 [0011] For example, in the case of using lithium-exchanged faujastic type zeolite as a crystalline zeolite for separating
- oir by PSA method, since the amount of nitrogen adsorbed is large, it is necessary to adsorb and desorb a larger emount of nitrogen as compared with the case of using a crystalline zeolite exchanged with e.g. coldium, and unless the diffusion race of nitrogen during adsorption and description is adequately improved, the adequate performance of the adsorbert can not be obtained. Further, in operating PSA, if the time for the adsorption and decorption step shortens, adsorption and description of nitrogen in a short period of time is required, and an adequate performance can not be obtained unless an adsorbent is used wherein the diffusion rate during adsorption and desorption is improved. Fur-
- ther, in order not to deteriorate the performance of the adsorbent itself, it is required to make the water content in the adsorbent as small as possible. [0012] It is an object of the present invention to provide an adsorbent for separating gases which is excellent in the
- as diffusion rate of the adsorbable component during description, and which has macropores having an average pore dismeter suitable for conditions for describing the advocable commonent in order to revise the electric power consumption by PSA appearatus, for separating gas mixture, particularly for separating gases by PSA method. It is also the object of the present invention to provide an adsorbent for separating passes which has macropores adventageous for adcorption and description of the adsorbable component, and at the same time which is excellent in strength properties
- represented by crush strength, and which has a small water content. Further, the present invention is to provide a process for easily producing such an adsorbent for separating cases. [0013] These objects have been achieved by the surprising finding that, due to studies on the macropore structure of the adsorbent for separating gases and the diffusion of the adsorbable component in the macropores, an adsorbent for
- soparating gases which comprises a binder and a crystalline low-sitics faujusite type zeolite with a SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub> notes ss ratio of from 1.9 to 2.1, wherein the average diameter of the macropores is equal to or larger than the mean free path of an adsorbable component when decorbing the adsorbable component, and at least 70% of the total volume of the macropores is occupied by macropores having a diameter equal to or larger than the mean free path of the adsorbable component, is excellent in the adsorption and description performance of the adsorbable component, particularly in the diffusion rate of the adsorbable component during description. Further, they have found that the detired adsorbert for
- ≪ separating gases can be obtained by adding water to e crystalline low-sitios fausasite type zeolite with a SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub>. molar ratio of from 1.9 to 2.1, and from 5 to 30 parts by weight of a binder based on 100 parts by weight of the crystalline low-silica faulabite type zeofite as dried, so that the bulk density is from 0.6 to 1.0 ke//, followed by kneeding, accomerating, calcinating, ion-exchanging and activating.
- [0014] Now, the present invention will be described in further detail.
- 45 [0015] The adsorbent for separating gases of the present invention is an adsorbent for separating gases which comprises a binder and a crystalline low-silice faujasts type zeolite with a SiO2/AI2O3 moler ratio of from 1.9 to 2.1, wherein the average diameter of the macropores is equal to or larger than the mean free path of an adsorbable component when describing the adsorbable component from the adsorbant, and at least 70% of the total volume of the macropores is occupied by macropores having a diameter equal to or larger than the mean free path of the adsorbable component. so 199161. For example, when superating gases by PSA method wherein an adsorbing step and a desorbing step are
- receased, pressure is exerted on the adsorbent during the adsorbing step and molecular diffusion mainly takes place wherein molecules diffuse while they collide against one another in the macropores of the adsorbent. However, the descriting step is conducted under reduced pressure, and the mean free path of the adsorbable component toricle to be large, and in the macropores of the adsorbert, collision of molecules against the well of macropores frequently takes ss place, in addition to molecular movement by the molecular diffusion. Accordingly, the diffusion resistance in the adsorb-
- ant during the describing step is larger than during the adsorbing step. Therefore, in the case where the average pore diameter of the micropores is smaller than the mean free path of the adsorbable component when describing the adsorbable component, or in the case where less than 70% of the total volume of the macropores is occurried by macro-

- pores having a diameter equal to or larger than the mean tree path of the adsorbable component, the diffusion resistence in the inside of the macropores of the adsorbable component is large, whereby the edicquate performance of the adorbable to sequerating assects can not be detailed.
- (0017) Further, it is preferred that the adoption for repositing gases also a total mecropror volume of at least 0.25 cold and a prox surface area of the less tot of mile. Linethy to major part of the macroprore of the exchange to depend on the expensiving gases is coupled by prote having a prox destriet of at least 1.000 Å. However, if macroprose are formed by such prox burking a proteinly large distriet value, the couple whereous the surface is the macroprose are formed by incorporating porce, burking a relative type affected value, the couple couple is made to that the couple of the couple o
- to total volume of macropores. Further, such perce having a relatively smell disenser-can serve as paths for gazas, whereby the dilation resistance of the setsorable component daving adeoption can be made small, as the mean free path of the adsortable component is small during the adsorbing step wherein the pressure is high.
- (0018). The mesoposes in the adoptive to asserting passes of the present invention are porce healing a pore distinion within a range of from 80 Å to 200 ym, as measured by the method of mesopre previously within a present any perror of from 1 to 50,000 pc. The everage pore dismeter of the macropores can be obtained from relation between the pore dismeter obtained by the method of mercury penetration and the pore volume (pore dismeter distribution curve), as a dismeter at 80% of the total pore volume (modified intenser) or a distinative when the gradatest of the pore dismeter distination.
- tribution curve is regularum (model dementa).

  [0019] The mean free pain of the actionable component is an average distance that molecules in the adsorbable component more culting the successive collisions. The mean free pain can be calculated from the pressure and the temporature at the stope of describing the adsorbable component from the adsorbabl ty PSA method, reterring to AL BRRYP VERSIOLA CHARMENT (from the folion 3,1924 to the PDSPF AL AB PRETY MEAN (ACCIVITIE) Fully.
- temperature at the stope of describing the adsorbable component from the adsorbart by PSA method, reterring to ALBERTY PHYSIQAL CHIMBITY (bourth design), 3129-14, by ROSERT A. ALBERTY, HAGANQL-GLYUTU PLUSHING CO., LTD. (DDD) The acceler of the adsorbart for separating gases of the present invention is a systalline low-sitica faulastic
- 21 type anothe with a BCA/M<sub>2</sub>O, moler site of them 1 5 to 2.1 flwerienistic retirest to as LSX acidits. The SCA/M<sub>2</sub>O, moler site of the LSX acidits is theoretically 2.9. However, within considering errors in measurement by the chainstal composition washless. It is evident that the LSX acidits believing a composition with a SCA/M<sub>2</sub>O, moler state of term 1.9 to 2.1 is within the unage of the present invention. Many methods for synthetizing a LSX receive with a SCA/M<sub>2</sub>O, most reason of term 1.9 to 2.1 have been disclosed. For example, it is possible to perfective the LSX acidits by a method of-colored.
- or closed in JP-9-5-25527.

  [0021] In the case of using the LSX secilite for an adsorbent for experiating gases, the higher the crystal purity of the LSX secilite for more excellent the esparating efficiency, and the crystal purity of the LSX secilite to preferably at least 50%. Measurement of the crystal purity of the LSX secilite can be carried out, for example, by an X-ray powder diffraction mellioff. by measurism the amount of one advanced for measurement amount of one advanced to the resource of the property of the crystal purity of the LSX secilite can be carried only for example.
- 55 method: [2022] As the bindor in the form of fibers to be used for the adsorbent for separating geas of the present invention, a bindor of needle crystals in the form of fibers is preferred to form managepose of the present invention, which commiss septicities day or strapdylip day, and which seeks, as a binder, among LBX zeotile particles in the adsorbent. Such days may be used allow or ea as misus-or be one or more of them.
- 40 [2023] If a binder in this form of plates is used, there is a possibility that the binder inhabits diffusion of the adsorbable component due to the shape. Further, weler is not quickly removed during calcinating after forming, whereby there is no
- possibility that crystal of the LSX souths is identroyed.

  [0024] The accordant representing passes of the present invention is preferably in the form of beads, and the shape
  [0024] The accordant representing passes of the present invention is not pranticularly limited on long as it has the characteristic of the advocation for separating gases of the present invention.

  In such may, for example, to product or eligibility. In credit or adoptively obtain the desired posturance of the
- udsorbant for soperating glaser, the dismeter is preferably from 0.5 to 5 mm, considering the size of the apparatus to be pucked, the present of copy in the peaked bed or the diffusion resistance in the inside of the applicaments. [0225] With regard to the compositional ratio of the LSX sociite to the birder in the adsorbant for separating gases of
- the present invention, considering the macropore structure and strength properties of the agglomerate, the compositional ratio of the LSX zeolite is usually preferably from 5 to 30 parts by weight. (9025) It is preferred that the water content of the absolutent for seperating gates is as small as possible, and the
- [duce] It is presented that the water content or the associated for experiming paying a part and processing, and the adoptions that which was the content of at most 0.5 wife, particularly all most 0.5 wife, shows a satisfactory exiscoprion performance and is preferably used.

  [00027] Now, the process for producing the adsorbent for separating gases of the present invention will be explained.
- 26 (2003) The process comprises existing value to a cytabilinit low-sizes business be paracillar with a SO<sub>2</sub>(A)<sub>2</sub>(A)<sub>2</sub> molar ratio of from 1.9 to 2.1, and store in 5.0 paratity by explicit of a binarial based on 100 parity by which of the caryballar cell calculate sizes the paracillar sizes of the calculate sizes the paracillar sizes of the calculate sizes of

[0029] The process for producing the adsorbest for separating passes of the present invertible comprises a step of acting a synthetized USX zeefile powder, a Birder and water followed by hisselfing, a sign of applications hiswards product, a step of dying and actionising the applications, a step of low-sucharaging the ballow degillomestic and a step of activating the low-schinged product by charatening. These steps will be explained below.

# Kneading step

(2009) The matrix greated for the sprinkers LSX solid provider to be used for the substitute for separating passes of hispersent investigation in a (MLV) by self-Loss symmetrized by the sprinker investigation in a (MLV) by self-Loss symmetrized by the sprinker investigation in the sprinker are mixed and insected visible codes for sometime for mixed and insected visible codes for sometime for mixed and insected visible codes for the bulk diversity in the sprinker solid providers and advantage for sprinkers and the first first and 5 self-logic throughout the sprinkers and the sprinke

(9052) In order to brinn the dissent amazopores is use produced in order to brinn the dissent amazopores, loop in plus allocation capacity and make the physical strength of the adoctornt high. The amount of the brinder adold is preferably within a range of them 5 to 30 parts by weight of the (Na.7) pp. 61.25 across. It is amount of the brinder is less than 5 parts by weight, the proportion of the LSX scale increases, each other amount of the brinder is less than 5 parts by weight, the proportion of the LSX scale increases, each other allocation amount of the brinder is less than 5 parts by weight, the proportion of the LSX scale increases, each other allocation amount of the brinder is less than 5 parts.

9 the amount of the bible blanch of the support of the support

[0033] The amount of water added when kneading the LSX zerolite powder and the bindor, varies depending upon the properties of the LSX zerolite powder and the binder as the starting materials, or upon the proporties of them. However, as the total amount of water added is preferably within a range of from 60 to 65 parts by weight based on 100 parts by weight of the LSX zerolite powder.

[0034] Further, as an additive besides water, an additive such as carboxymethylcellulese or a polyvinyl allochol may be added.

# so Applomerating step

(2003) The knotted inducts having a bulk density of them 0.0 to 1.0 by if a againmented as blows. It is preferred him to the induction of a collegement by the date of planting manufacts, including production method, the brinds existed is disposed united to the collegement of the

42 ing a High aghirotich, It is aposition to gramulate the agglomerated spherical product by a known method such as using MAPUMERIZER forming apparatus, to make the surface of the agglomerate amount. (2008) The agglomerate in the extraction of the agglomerate in the agglomerate to beart, it is possible to obtain the desired macropera structure, by using cates output/plottables or a polyveyl absolve that her known as beinging adgl, and the agglomerate that agglores structure, by using cates output/plottables or a polyveyl absolve that her known as beinging adgl, and the agglores agglores agglores.

66 agglomerate in the shape of a pellet can be used as an adsorbent for separating gases. (0037) That dismetter of beads which is agglomerated and granulated can be changed depending upon applications, and it is costable to classify the size by using a.g., a sieve.

### Calcination step

[0098] The apploments thus obtained is dried and calcineted, and the binder added was calcineted. As a method of driving and calcineting, a comentional method can be employed, and a girl and air or, a multi-braneau, a rollary life or a bits forman employee. On the imperation of calcinating may be earlier brander in calcineting and the service and section of the production of the calcineting and section of the calcineting as a conducted at a transportation of from 400 to 700°C.

[0039] Further, astainsted egglomerate is ocaled and may be humidified so that the water content is from about 20 to about 30%. Although humidifying operation is not assential, it is effective to prevent breakings such as cracking of the aggloments resulting from a sudden hesting due to well-read-pring when contenting with the on-exchance but on

during ion-exchanging in the next step, and it is also effective to evacuate the acts about gas such as nitrogen from inside of the aggiomerate, and to make the cilibasion with the ion-exchange solution effective.

#### lon-exchanging step

[0040] The aggioments formed and calchinated in the above steps, is consisted with an ion-exchange existion having caches such as fifthin, potentiars, noticulars, strontisms or brisms, to conduct or exchanging, the type of calchinate selected depending upon the gas absorbed. For exemple, in the case where introgen in the air is actionated to expense the sit, tilbure cancin is suitable. The compound to be used for for exempling is not particularly intended so long as it the sit. Elibure cancin is suitable. The compound to be used for for exempling is not particularly intended so long as it.

10 can be in a form of an aqueous solution, and a chloride, a nitrate, a suifate or a carbonate may, for example, be preferably used.
100411 As a method for lon-exchanging, batch method or column flow method is usually employed. The batch method or longer than the longer than

[D843]. As a method for low-sechanging, batch method or column flow method is usually employed. The batch method is usually the column flow method is usually the programment of the programment of the substance of the programment of the progr

the column flow method is preferred.

[9042] The temperature during ion-exchanging is preferably as high as possible in order to improve the rate and effi-

clency of lon-exchange. It is usually from 50 to 100°C.

(043) The concentration of ion-exchange solution to be used is generally from about 1 to about 4N, considering the pro-exchange rate. The increadurate solution is oreferably sileatine so that LSX zeedits crystal is not destroyed during

ion-exchanging, and it is generally adjusted to have a pit (hydrogen ion concentration) of from 9 to 12, by adding a hydroxide or the file. [0044] After ion-exchanged as mentioned above, the applicaments it taken cut from the ion-exchange solution, weeknot with water or his water, and ride at at temperature or usually them about 50 to about 10°C.

#### Activating step

(2004). The againment from the recharged in arthritish to remove water, and the destined accorded to separating gaster can be obtained. The purpose of carbolisms in to remove whether in the againment of their is removed under an extensive present on the control of the purpose of carbolisms in the present present and control of sectionary in particularly their internoval under a depotement in the case where the againment of scholations may be any condicion whether the removalement and approximate in the case where the againment of scholations are passed to a provide, and the present the removalement of the adjustment of the scholation of the SSX and/or in the scholation of the scholation

27 [0046] The addictions for expectating glass obtained in the above steps, is used to actorb and expense the mixed gas by addicting the addicatable component in the mixed gas for expension and concentration. For example, it is used to recover concentrated congrue pasts by selectively addicting altergain in the air. The process of concentrating and recovering copying as in the air by PSN method is operated by successive steps of an advorting step wherein the air concentration.

is consisted with the product box and management associated by the control of the product box of box of box of the product box of bo

or cooled in order to obtain an adequate performance of the assorbent. However, it is usually from about 15 to about 55°C.

50°C.

50°C

range of from 760 for to 1,500 for.

[D048] The lower pressure in the regeneration step is preferred since more nitropen can be desorbed. Considering the lossifing to the viscuim pump, the regeneration pressure may be within a range of from 100 flor to 400 for.

[D049] In the pressure-reginging step, as the concentrated conjugo gas obtained in the advancing step is used, if

or Tegulined pressure is high, this amount of the concentrated copying data taken out as a product gas accesses. Further, this advocating gas starts in the state where the regarding pressure is line, was pressure is except to the except former with an intercept in the state where the regarding pressure is low, as pressure is exerted to the exit, there are fearer than introgen in the act advocated on the acknowled must be transpired to the pressure may be regained by ordering the control of the pushed both. In pressure may be regained by ordering the

- concentrated exygen gas back to the packed bed counter-currently to the air for the initial period of about 1 to about 5 seconds after the east-ording step started. The return pressure may be within a range of from 400 Terr to 800 (1905). With regard to the automother for repetinging packed of the present invention, the paveage diameter of the micro-
- pore is equal to or larger than the mean the path of the astochable component when described per advocable component when described per advocable component from the advocable component from the advocable component from the advocable component from the advocable component. Therefore, the distillation that of the advocable component in the advoc
- no pore d'arméter are suisably incorporated among macropores in addition to pores having a restrively lenge pore d'arméter (0051). The addrether to resperating gasses of the present invention is more effective for separating the sair by PDA method. The ritter, the amount stating not used the reconsciration conjugate, ser leigh which the association by PSA method is conducted, end it is possible to reduce the electric power consumption rate when operating the PSA construct.
- ing the PSA apparatus.
- EXAMPLES

[0052] Now, the present invention will be described in further detail with reference to Examples. However, it should be understood that the present invention is by no meane restricted to such specific Examples. Each evaluation is based on following methods.

# (1) Average pore diameter, pore volume and pore surface area of macropores.

[0003]. By using netwary proximeter (produced by MICROMERTICS, type: PORE SIZER 9810), activated activates of various measured within a pressure respect from 1 to 3,000 for sident power dismers and from 60 A to 200-mil. From the relation between power dismers and previous obtained by measurement (pow dismers distribution curve), the surveys power distinsted or sident power distribution curve), the surveys power distinsted or the distribution curve) and of the surveys power distribution curve). The surveys power distribution curve (and individually or a distance when the grade of the power distribution curve is maximum (modal dismetler), in the prosent Ediment when the grade of the power distribution curve is maximum (modal dismetler).

# (2) Amount of nitrogen edsorbed

- (DSIG). For measurement in Exemples 1 to 15 and Companible Exemples 1 to 12, garametric method was employed by using an electric believe (DAIR\*) bodge). Also by printing restatement, the active draw assistant for so as at temperature of 50°C under seasons of at most 10°C for. The active global subpretative was feel at 10°C, 0°C or 20°C, rimprogram grows extractable, and the weight was researched in the address described in the afficient settle or Exclusion active to active amount of a most of the amount active and the afficient season of the amount of the address are active and the amount of the address and the activity of the amount accorded (with Notice). Each amount of inhops addressed in the address according to the activity of the acti
- (0055) For measurement in Examples 15 to 18 and Comparative Example 13, voluntaritio method was employed by or using BELSORP 255A (BEL JAPAN, NO.). As the preliminary treatment, this autobative are subjected to dealing treatment for 2 bound at room temperature (about 25°C) under vacuum of at most 10° Tor. The addrespition temperature valloopt at 25°C, and this amount addressed was measured to about 800 Tor (unit: No.)). Each amount of infrequent addressed in Exemples and Comparative Example is an executed value under 700 Tor.

#### 47 (3) Air separation test by PSA method

- [0056] By using a testing device for air-separation performance as illustrated in Figure 1, air separation test was conducted as follows. About 2 Left elasterbant for expansing gases was packed in adsorption bed (13) and (14). While on adsorption the (13) was in the adsorption shot (13
- so drafting bad (3), the pressure was reduced by a pressure reducing valve (6) to 5 cm .5 to 0.6 legicam<sup>2</sup>Q and celeroid valves (5) and (7) were feet open to flow the six filteruph the adsorption bad (six interpretature 20°C). The obtained concentrated organic pass were reserved in a product fault(17), and the amount of the concentration organic pass belief or was adjusted by a mass flow meter (18). The pressure at the final of the adsorbing step was constant at 1.4 arth. While the adsorption bod (19) was in the regineration despit the soldered valves (2) and (7) were obecast, an advanctive to the soldered valves (2).
- ss was kept open, and pressure in the inside of the edecoption bed was reduced by a vacuum pump (20). The fixed precurs at the fixed of the reoperation steps was content at £20 Tim. While an adoption bed (13) was in the pressureregaining step, the ademoid valve (6) was closed, an oblenoid valve (8) was kept open, end the pressure in the linkled of the addorpion bed was respined by the condensited oxygen one or the product start (17). The pressure at the final

of the pressure-registring slap was constant at 500 Torr. The procurs was measured by a pressure gauge (16) fitter and addropping the CV), a pressure gauge (16) see sured). Then the absorbing slap was considered in the pressure-registred addropping the CV), and pressure-registred addropping the CV), and these slaps were successively repeated. The time for the adopting stap, the registration step and the pressure-registring stap were 1 minute, 30 seconds and 50 seconds; respectively. The consistency and the state of the consistency and the consistency are consistency as the consistency are consistency as the consistency are consistent as

- or the scale not of whose ware controlled by a sequence.
  [DORY] The scale not on some caso are conducted in an adaptorption bad (14), in order to confinencely takes out the concentrated congert gats, the subcorption bad (14) was in the regeneration stop and the pressure-registring stop while the adorption bad (14) was in this adorbting test pan of the adorption bad (14) was in the adorbting test with the adorbting test with the adorbting test pan of the adorption bad (14) was in the adorbting test pan it are adorbting to the subcorption of the adorbting test pans th
- wat is the regeneration drap and the pressure-regalning step.

  [2003] The concentration of the concentrated oxygen gus was measured by an oxygen meer (10) after the vestice became containt, and the preside serrount of the concentrated oxygen gas flowing (the chartler related to as serrount of oxygen) was obtained from the value measured by an integrating flow meter (20). And the amount of advantated gas evanuated or the responsable of the things of the chartler of the chart
- ovicusinal from the visuum pump (20) ulving the regeneration step (heritarities referred to as amount of similarities) goal was children direct the value measured to part hereoffee (20). Each mencent of gas was measured to an integrating flow merit (20). Each mencent of gas was measured at 20°C.

  [10059] The air-separation performance of the adoctobest was represented by the amount of oxygen at a concentration
- of 93% and the ratio of the concentrated expensions as a concentration of 93% recovered from the air (hereinstate referred to as recovery ratio). The air expension test was conducted at a temperature of the adsorption bad of 0°C and 25°C.
- 20 [0060] The amount of copyen was obtained by calculating the value measured by the integrating flow meter to the normal state, and represented as an amount of copyen flowing in 1 hour per log of the adsorbert as direct. The unit is NU[fig -rh]. The recovery ratios was autostated from the toliving formula.

Recovery ratio = ((smount of cxygen) x 0.93)/((amount of the air supplied) x 0.209) x 100 (%)

Amount of the air supplied - (amount of oxygen) + (amount of exhausted gas)

(4) Mean free path of nitrogen when nitrogen is described

- 200511 The mean rise paint of introgen (2) was calculated from the blowing formula by unlong the temperature of the actions/plen bod (7) and the first procure at the regeneration stop (P) which are conditions for the air separation test as minimized above, in the case of separating the air by PAI method, the expressable of the advoctment raises by hearing in the advocting is not the temperature of the advocting stop, thus leading to fluctuation of temperature of the advocting stop, thus leadings to fluctuation of temperature. The residence was recommended to the control of the procedure of the control of the procedure of the control of the control of the procedure.
- 55 temperature of the adsorbent, the temperature of the adsorption bed (ambient environment) or the temperature of gas introduced may be employed. In the present Example, the temperature of the adsorption bed was employed.

 $\lambda = \kappa T/(\sqrt{2})\kappa P\sigma^2$ 

φ σ : Molecular diameter of nitrogen 3.681 × 10<sup>-10</sup> (m)

π: Boltzmann's constant 1.3807 × 10<sup>-23</sup> (J/K)

[0062] The mean free path of nitrogen under a pressure of 250 Terr, was 2,052 (Å) or 1,890 (Å) at a temperature of 25°C or 0°C, respectively.

(5) Gruth strength

[0003] 25 activated agglomerates were measured by a hardness mitter (produced by KYA SEISAKUSYO LTD., type: (HT-60). The measurement was concluded in the method such that the agglomerate was loaded with an infant hardness as diameter of 5 mm at a constant rate of 1 mm/beor. The amount loaded when the agglomerate broke was taken as the onus framenoff cost.

(6) Attrition rate

50041. The driftion nate was coloulated in accordance with a measuring method as sipulated in JIS K-1-464 (1982), Namely, the orderided agreements as a text transpla was preliminarily high for 16 hours in a desposalor as the empretaire of 25°C at a relative humidity of 80% to reach equilibrium. About 70 g of the test sample was subjected to steving for 3 minutes by unique a screen having deverse 680 ym and 435 ym and a page (page 400 ym 70 NYO) SCREEN COL LTD.

type: IIIS Z-8801). Then, S0 g of the remaining test sample obtained by the severing ter 3 minutes, was proctary manured and put on the above screen when daying deposits removed therefrom. At the same time, for We 10-yes opposit, or put in the screen, followed by shalling for 15 minutes. The amount dropped to the pan was taken as X g, and the attrition rate was obtained by the following termula.

### Attrition rate (w/%) = (X/50) x 100

#### (7) Bulk density

10 (0065) The bulk density was measured in accordance with a method using an apparent density appearatus as stipulated in JS K-GREZ. The invasided mixture was upon but in a polyethylence cup of V in (WI) so that the cup was heaped with the invasided product, the invasided mixture was caread of the gilled product, the invaside printed was expressed to the unit of 0.1 g, and the bulk density was calculated by the following remains:

#### Bulk density (kg/t) = (W2-W1)/V

## (8) Water content

20 (006) The warde content was measured in eccordance with the test method by coulomatic literation as eligibated in JIS K 0068, by using Kerl Facher microture mater (produced by MITSUBSH CHEMICAL CORPORATION, moisture measurement mater: CA 05 type, electric barrace 3/4/21 type). The electric barrace was set to 400°C. From electric to be about 500 mg of the achiested lest cample was precisely measured quietly, the test example was put in the serropic boat in the selectric harmace, and washin was expected under folking day in from an 430 m/limit. The water content was

boat in the electric furnace, and water was exported under flowing dry nitrogen at 300 mi/min. The water content was obtained by the following formula from the amount of the test sample (S, unit; g) and the amount of water obtained from contenents fination (S, unit; sa).

#### 41 FYAMPLE 1

[0067] Synthesis of LSX zeolte was conducted by a known method. To a stainless container for reaction having an Inferned volume of 20 4,3,883 g of squeebus sodium atuminate solution (N<sub>NC</sub>O = 20.0 w/th, N<sub>C</sub>O<sub>2</sub> = 22.5 w/th), 7,833 g of water, 2001 myldragetim hydroxide (purity 59%) and 1,485 g of special practic chemical protection hydroxide (purity 59%) are 1.485 g of special practic chemical protection hydroxide (purity 59%) were

- 27 put and cooled under stirring at 80 rpm (solution g; SO). To a polyethylene container having an internal volume of 10 £7.750 g of acquous souther alterial solution(s)<sub>2</sub>O = 3 a wm, SO<sub>2</sub> = 12.6 w(3)<sub>2</sub> and 1,176 g of water where put and cooled (solution(g) 10°O). This solution have put into the solution gover a polyethylene under stirting. The solution after mixed water transporters. Stirring was look about 20 includes after completion of putting solution it. Then, the temperature of water both they are lasted 95°C. As one as the solution after and closely stirring was stooded the stirring of the solution of
- 4: ring pastide was taken out, and aging was conducted at a temperature of 80°C for 48 hours. (2008) Then, the temperature of water both was naised to 70°C, and exystalization was conducted for 20 hours. The obtained crystal was subjected to filtration, adequately washed with pure water, and disco overrigit at a temperature of 50°C. As a result of X-ray analysis, the structure of the obtained erystal powder was faultation completion, and had a purity of all seasons 95%. Further, or a result of ICP entition analysis, the composition of the obtained crystale product was fault and crystal product analysis.
- 7.28/kg.O 0.28/kg.O M<sub>cO.</sub> > 2.08(b<sub>c</sub>, and it is confirmed that the crystal powder was LSX zeotine. (1008)
   The LSX zeotine powder and 20 parts by weight of attapulgate cuty based on 100 parts by weight of the LSX zeotine powder were mixed and visuaded by MIXMULEER MIXING MIXMUNES (produced by SINTOK/COI).
   I.T. by the SINTOK/COI, and the size of the LSX zeotine powder was a fixed powder. Firstly, 65 parts by weight of whate based on 100 parts by weight of the LSX zeotine powder was added, Informed by forendard. The bud known by of the College of the SINTOK/COI, and the size of the
- 50 (2079) The invested product was sirred and terms of the basels having a distinster of from 1.2 to 2.0 mm, by blade agistion grounder H-PINGHER. USARS (Produced by MILIN MINIO COUNTY, ICE, type, PIN-FIN, which was agranulated by using MARILMERIZEF EXTRIDER (produced by FILIN PILINAL CO., LTD., type, C-1000), filinated the distinct of the produced by FILINAL CO., LTD., type, C-1000), filinated of 500 County of the PIN-FIN MILINAL CO. LTD., type, C-1000, filinated and Section 1.0 of the County of the PIN-FIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the County of the PIN-FIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the County of the PIN-FIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the County of the PIN-FIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the County of the PIN-FIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the County of the PIN-FIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the County of the PIN-FIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the County of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the County of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of the PIN MILINAL CO., LTD., type, C-1000, filinated and Section 1.0 of t
- [0071] The egglomerate was packed in a column of 70 mm/2 × 700 mm (length), and an equation solution having lithium chloride adjusted to a contentiation of 1 mm/2 was flowed at a temperature of 80°C to contact lithium interactivings. Then, the agglomerate pecked in the column was adequately washed with pure water, and taken out to

the column, followed by drying for 16 hours at temperature of 40°C.

[6072] Then, the agglowments was subjected to activation treatment by using a harizontal type tube flumace (produced by ADVANTEQ) for 1 hours as impresenture of 50°C under flowing the air, and wrapped without cooling. The everage

cy AUVN-1 LtD, lot 1 nour as a temperature of 500°C under flowing the air, and watpool without cooling. The average pero distinate rand the pore obtained in the destinate distorted for expensing gases were measured. This amount of through endoubted and the air-expension performance were measured at an edeoption temperature of the above the second of the cooling of the cooling

#### EXAMPLE 2

[0073] The same operations as in Example 1 were conducted, except that the bulk density of the kneeded product was 0.90 kg/l. The results of measuring the obtained adsorbent for experating gases are shown in Table 1.

#### EXAMPLE 3

[0074] The same operations as in Example 1 were conducted, except that sepicitie city was used as a binder. The results of measuring the obtained adsorbent for separating gases are shown in Table 1.

# EXAMPLES 4, 5 and 6

[0075] By using the same adsorbent for separating gases used in Examples 1, 2 or 3, respectively, the amount of nitrogen absorbed and the air-separation performance were massared at an adsorption temperature of 0°C. The resurts of measuring the obtained absorbent for separating gases are shown in Tebio 2.

#### 25 EXAMPLE 7

[0079]. The same operations as in Exemple 1 were constanted, except that it 5 parts by weight of exploits clay was used as a binary. The results of measuring the pero volume, the pore currice area, the crush strength, the striction rate area of the embass of the embas

# EXAMPLES 8 to 15

[0077] The same operations as in Exemple 7 were conducted, except that the type and the amount added of the binder were changed as aboven in Table 5, to oppose an advantant to expensing gause. The balk density of each invaded product measured by the above method, was within a range of from 2014 to 0.0.7 (e/c. The pore volume, the pore curked area, the crush strength, the attribute mate and the amount of introduce selections (acknowledges temperatures: -10°C) of the duthinal advantant for expensing passes are shown in Table 3.

#### 40 EXAMPLE 16

[0073] The same operations as in Example 1 were conducted to stiffned the advantent for operating group. Bettern the additionate was regional, as it states not from the factor to the conduction of the conduction and the states are stated in a glass both and the states of the states of the state of the states of the states

# EXAMPLE 17

[0079] The same operations as in Example 16 were conducted except that the adsorbent before wrapped was cooled to 350°C. The measurement results are shown in Table 5.

# **EXAMPLE 18**

[080] The same operations as in Example 16 were conducted except that the adsorbent before wrapped was cooled to 300°C. The neasurement results are shown in Table 5.

#### COMPARATIVE EXAMPLE 1

[0081] The same operations as in Example 1 were conducted except that the built denotity of the adsorbest after impacted and mixed west 1.8 kg/l, and forming was conducted by using MARIAMERIZER (produced by FLUI PAUDAL O

# COMPARATIVE EXAMPLE 2

[00022] This same operations as in Example 1 were conducted except that the bulk density of the advoctors after insteaded and inheal was 12 kg/l, and training was conclused by using MARIMERIZER (Produced by PLI) PRUINAL CO., LTD, type: Q-1000) alone. The measurement results of the obtained advoctors for expending gases are shown in Table 1.

### 15 COMPARATIVE EXAMPLE 3

[0083] The same operations as in Example 1 were conducted except that the bulk density of the adsorbent after kneeded and mixed wite 1.2 kg/L. The measurement results of the obtained adsorbent for separating gazes are shown in their 1.

# COMPARATIVE EXAMPLES 4, 5 and 6

[0884] By using the same adsorbent for separating gases used in Comparative Exemples 1 or 2, the amount of the manager actions during the separation performance were measured at an adsorption temperature of 0°C. The measurement results of the obtained adsorbent to separating gases were although Table 2.

#### COMPARATIVE EXAMPLES 7 to 12

[0:085] The same operations as in Examples 7 were conducted except that the type and the amount added of the 50 bindle were changed as shown in Table 4, to proper an adaptive for exposating ages. The build density of each kneeded proton was measured by the above method, and found to be within a range of term 0.56 to 0.50 kg/l. The results of measuring the poor volume, the prior surface area and the amount of introgen adviced (adaptivition temperature. 10°C) of the obtained additionable for securities or uses were shown in Table 4.

#### Ø COMPARATIVE EXAMPLE 13

[0036] The same operations as in Example 16 were conclusted except that the adsorbent before wrapped was cooled to 200°C. The measurement results are shown in Table 5.

- [0087] With regard to each adsorbant for reparating gaines shown in Examples 1 to 6, the evenage point diameter of 5 menaproces was larger than the mean free path of introgen under 250 Torr under which introgen was described during the bit-expiration by IPS4 process, and at least IPS6 of the total pore volume was occupied by the pores having pore diameter of depail to or larger than the mean tree part, and the diffusivity in the mercupour under condition of 250 Torr was high. Therefore, the air-experization portnames of the adoctories to respecting posses was occulant.
- [0088]. With repart to leach individual for expending passes of Companion Example 1 or 4, the average por dismates of the harcecopers was emailed that the ment here paid or inference used 250 for under what indepen was actioned using the all-operation by PSP process, and less than 7% of this total para volume was occupied by the porces benefit passes above the contract of the high process and the passes of the three passes of the passes of the passes of the passes of the three passes of th
- Section of the companion of the compa
- [0089] Further, by comparing Examples 7 to 18 and Comparative Examples 7 to 12, it was found that the adoptions of obstacle in Examples had a higher pore votine, pore surface area and amount of nitrogen adoption, and were more exhallent as an articular the secretain nation.
- [0090] By comparing Examples 16 to 18 and Comparative Example 13, it was found that the adsorbents obtained in Examples having a water content of at most 0.8 wt%, had a higher amount of nitrogen adsorbed, and were thus prefer-

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## able as an adsorbent for separating gases

adsorbents at an adsorption temperature of 25°C Table 1 Air-separation performance of

Amount of Recovery 61.5 ratio Air-separation performance 103.9 103.2 Amount of nitrogen adsorbed (25°C) (Ncc/g) 21.8 21.4 Proportion of pores having a diameter of at least 2052 A volume (%) 81.5 71.9 Pore volume (cc/g) 0.27 0.32 pore diameter (Å) Examples | Type of clay Average 5000 4300 Attapulgite Attapulgite œ

61.2	Air-separation performance Amount of Recovery oxygen ratio	æ	58.2	100	70.1	57.7	
102.6	Amount of Air-separation initrogen performance adsorbed Amount of Reco	(ML/kgh)	95.1	99 2		95.5	
23.2	Amount of nitrogen adsorbed (25°C)	(B/cc/d)	21.2	22.0		22.4	
1.20	Proportion of Amount pores having a nitrog diameter of at adsorb hans 2052 A in (25°C)	(4) 2000	0.82	65.6		0.09	
0000	9.0	90.0	9	0.32		0.30	
200	Average Pore pore diameter (co/g)	1490		2890		2700	
	Comparative Type of clay Average Pore poxemples diameter (co/g	Attabuloite		Attapulgite	National aday	occapation of	
	Comparative	1		2			

000

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duning P	S A A E		Performance of the control of the	Pore volume (cc/g) 0.27 0.32		Amount of Mitrogen Mitrogen Macorbed (0°C) (Mcc/g) 34.2 33.9	Secretarion temperature of 0°C	of 0°C Eion e Recovery ratio (4) 56.1 56.1
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tion	Recovery	6 62	3	: 3	3
Amount of Air-separation nitrogen performance	Amount of Recovery oxygen ratio	92.5	9 90	5 53	
Amount of nitrogen	Amount of nitrogen adsorbed (D°C) (Ncc/g)			34.9	
Proportion of pores baving a diameter of at	least 1880 A in volume (%)	32.0	68.8	63.3	
Pore volume (cc/g)		0.25	0.32	0.30	
Average pore diameter	(A)	1490	2890	2700	
Comparative Type of clay Average Examples Dore diameter		Attapulgite	Attapulgite	Attapulgite	
Comparative Examples		·	2	٥	

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Table 6

Example	s Type of clay used	Amount of clay used (parts by weight)	Pore volume (co/g)	Pore sur- tace area (m²/g)	Crush strength (kgf)	Attrition rate (%)	Amount of nitrogen adsorbed (- 10°C) (Noc/g
7	Sepíolite	15	0.31	21.9	1.5	0.1	32.0
8	Sepiolite	15	0.31	22.7	1.6	0.2	31.5
9	Sepidite	20	0.29	28.2	1.9	0.1	30.8
10	Sepicite	20	0.27	28.1	2.1	0.1	30,6
11	Sepicite	25	0.26	32.3	2.4	0.1	29.9
12	Attapulgite	15	0.30	22.0	1.3	0.3	31.0
13	Attapulgite	50	0.28	25.7	1.8	0.1	30.0
14	Attapulgite	20	0.28	26.1	1.9	0.1	30.1
15	Attapuigite	25	0.25	35.1	2.3	0.1	29.2

Table 4

30	Comparative Examples	Type of clay used	Amount of day used (parts by weight)	Pore volume (oo/g)	Pore surface area (m²/g)	Amount of nitro- gen adsorbed (- 10°C) (Noc/g)
	7	Kaolin	15	0.24	16.9	29.0
	8	Kaolin	20	0.23	19.8	28.1
35	9	Kaolin	25	0.21	21.1	27.9
~	10	Bentonite	15	0.23	16.5	26.7
	11	Bentonite	20	0.22	18.6	25.8
	12	Bentonite	25	0.22	19.7	24.3
40	The amount of clay	used (parts by we	eight) is based on 10	parts by weight o	of zeolite.	

Table 5

	Water content (wt%)	Amount of nitrogen (25°C) (Noc/g)
Example 18	0.30	20.8
Example 17	0.48	20.0
Example 18	0.71	18.2
Comparative Example 13	0.98	16.3

(0091). As mentioned above, with regard to the adsorbent for separating gases of the present invention, the average pere diameter of the macropores is larger than the mean free path of the adsorbable component when desorbing the

adsorbable component, and at least 70% of the total macropore volume is occupied by porce having a pore clameter of equal to or larger than the mean free each of the adsorbable component. Therefore the diffusion rate of the adsorbable able component in macropores during description under reduced procesure is tigh, and the utilization ratio of the artsorbent is high. Further, by suitably incorporating pores having a relatively small diameter, it is excellent in strength

- s properties. The adsorbent for separating gases of the present invention is more effectively used when it is explanded with lithium cations, and it is used for according the air by PSA method. Therefore, when senarating the pir by PSA method, the amount of the concentrated oxygen gas taken out and the recovery ratio are high, and it is possible to reduce the power consumption when operating the PSA apparatus. Further, according to the process of the present invention, it is possible to easily obtain an adsorbent for separating pases.
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#### Claims

- 1. An adsorbent for separating cases, which comorines a binder and a costalline low-silics faulosite type profite with a SiO<sub>2</sub>/Al<sub>2</sub>O<sub>3</sub> malar ratio of from 1.9 to 2.1, wherein the average gore diameter of the macropores is social to or larger than the mean free path of an adsorbable component when describing the adsorbable component from the adsorbent, and at least 70% of the total volume of the macropores is occupied by macropores having a diameter equal to or larger than the mean free path of the adsorbable component.
- 2. The adsorbent for separating gases according to Claim 1, wherein the macropores have a total volume of at least 0.25 colg and a pore surface area of at least 20 m2/p.
- 3. The adsorbert for expensing gases according to Claim 1 or 2, wherein the purity of the crystatine low-siles faujaette tyne zwolite is at least 90%.
- at 4. The adsorbent for separating gases according to any one of Claims 1 to 3, wherein the binder is sepicitie day entifor exaculate alov
  - 5. The adsorbert for secessing cases apporting to any one of Claims 1 to 4 which is in the form of breats
- 35 6. The adeorbent for separating gases eccording to any one of Claims 1 to 5, wherein the crystalline low-silica faulasite type zeolite is ion-exchanged with lithium cations.
  - 7. The adsorbent for secerating cases according to any one of Claims 1 to 6, which has a water content of at most 0.8 wt%.
- B. A process for producing an adsorbent for separating gases as defined in any one of Claims 1 to 7, which comprises adding water to a crystalline low-sitics faulacite type zeolite with a SiO<sub>2</sub>/Al<sub>2</sub>O<sub>2</sub> molest ratio of zeolite crystal of from 1.9 to 2.1, and from 5 to 30 parts by weight of a binder based on 100 parts by weight of the crystalling low-sitios
- faultisite type segite as direct, so that the bulk density is from 0.8 to 1.0 kg// followed by knowing, accommission. 9. The process for producing an adsorbent for separating gases according to Claim 8, wherein the binder is sepicitie clay ant/or attapulgite clay
- 45 10. The process for producing an adsorbent for separating cases according to Claim 8 or 9, wherein the adsorbent is formed into beads by blade agitator granulation.

calcinating, ion-exchanging and activating.

- 11. The process for producing an adsorbent for separating gases according to any one of Claims 8 to 10, which comprises ion-exchanging the calcinated applomerate with a solution having lithium cations.
- 12. A process for separating gases, which comprises contacting a mixed gas with a packed bad at the adsorbent for separating gases as defined in any one of Claims 1 to 7, to solectively adsorb at least one constituting gas in the mixed gas.
- ss 13. A process for separating nitrogen gas-oxygen gas, employing the process for separating gases as defined in Claim 12, wherein the gas is sir, and nitrogen gas is selectively adsorbed to recover oxygen cas.
  - 14. The process for separating nitrogen gas-axygen gas according to Ciaim 13, wherein nitrogen in the six is selectively

m

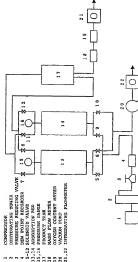
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# adsorbed by a pressure swing adsorption method.

- 15. The process for expansing shinging piece-organ pas according to Calin 14, which is operated by an also ofting speakershine in its ordated with the possible of and integrate is selectedly advantited for process processing organs from the coate of the passed bod; is represented in step wherein the contact of the size with the passed bod; is represented in step wherein the contact of the size with the passes are the size of the possible of the inducted bods on bottomed integrate and the volculate is card a pressure respairing step wherein the incide of the passes bottomed in these and the volculate is card a pressure respairing step wherein the incide of the passes body and the passes are respectively.
- 16. The process for separating ritrogen gas-cocygen gas according to Claim 15, wherein the adsorption pressure during the adsorbing step is within a range of from 750 Torr to 1520 Torr.
  - The process for separating nitrogen gas-oxygen gas according to Claim 16 or 16, wherein the recovery pressure in the regeneration step is within a range of from 100 Torr to 400 Torr.
  - 18. The process for separating nitrogen gas-oxygen gas according to any one of Claims 15 to 17, wherein the regained pressure during the pressure-returning step is within a range of from 400 Torn to 800 Torn.





SCHEMATIC DIAGRAM LILUSTRATING TESTING DEVICE FOR ALE-SEPARATION PERFORMANCE